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The ciliopathy-associated CPLANE proteins direct basal body recruitment of intraflagellar transport machinery

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Summary/Abstract:

Cilia use microtubule-based intraflagellar transport (IFT) to organize intercellular signaling. The ciliopathies are a spectrum of human disease resulting from defects in cilia structure or function. Mechanisms regulating assembly of ciliary multiprotein complexes and their transport to the base of cilia remain largely unknown. Combine proteomics, *in vivo* imaging, and genetic analysis of proteins linked to planar cell polarity (Inturned, Fuzzy, WDPCP), we identified and characterized a new genetic module, which we term CPLANE (ciliogenesis and planar polarity effector) and an extensive associated protein network. CPLANE proteins physically and functionally interact with the poorly understood ciliopathy protein Jbts17 at basal bodies, where they act to recruit a specific subset of IFT-A proteins. In the absence of CPLANE, defective IFT-A particles enter the axoneme, and IFT-B trafficking is severely perturbed. Accordingly, mutation of CPLANE genes elicits specific ciliopathy phenotypes in mouse models and is associated with novel ciliopathies in human patients.

Introduction:

The ciliopathies are a broad class of human diseases that share an etiology of defective cilia structure or function. These diseases span skeletal anomalies, craniofacial defects, cystic kidneys, blindness, obesity, and other presentations, highlighting the wide array of physiological functions that require components of the cilium^{1,2}.

Like all organelles, cilia are assembled and maintained by multi-protein machines. For example, the BBSome is a large complex involved in trafficking of ciliary membrane proteins³; the Nphp and Mks/B9 complexes assemble the ciliary transition zone, which controls access to the cilium^{4,5}; and dynein arms drive motile ciliary beating⁶. Likewise, the intraflagellar transport (IFT) system, which links cargos to microtubule motors for transport into and out of cilia, is comprised of two multi-protein complexes, IFT-A and IFT-B⁷⁻¹¹. IFT-A and IFT-B are frequently described as controlling retrograde and anterograde traffic, respectively. However, both anterograde kinesin motors and retrograde dynein motors can physically associate with IFT-A^{7,12-14}, and recent studies highlight the role of IFT-A in ciliary entry and anterograde traffic^{15,16}. IFT-A and IFT-B are each composed of a multi-protein “core” in addition to more loosely-bound “peripheral” components^{15,17-19}.

While recent papers have begun to define the interactions between IFT complexes and their cargoes (e.g. Ref. ^{20,21}), substantial questions remain concerning the mechanisms by which IFT proteins are recruited to the base of cilia and assembled into IFT trains. Indeed, ciliopathies can result from defects in cytosolic factors that facilitate dynein arm or BBSome transport and assembly²²⁻²⁸, yet little is known about similar factors that may act on IFT. For example, basal bodies in *Ttbc2* mutant mice fail to recruit certain subunits of the IFT-A and IFT-B complexes, but they also fail to remove CP110, a key initial step for ciliogenesis²⁹. Likewise, *Odf1*, *Cep83/Ccdc41*, and *C2cd3* are implicated in recruiting certain IFT-B subunits to the basal body, but the specificity of these proteins for IFT recruitment is hard to discern, as only a small subset of IFT proteins has been examined in these mutants and these proteins have pleiotropic roles in ciliogenesis³⁰⁻³².

Here, we combine proteomics, *in vivo* cell biology, mouse models and human genetics to characterize a novel mechanism governing basal body recruitment and assembly of IFT-A. This novel regulatory module is formed by specific protein-protein interactions among *Intu*, *Fuz*, and *Wdpcp*, well-conserved proteins that control planar cell polarity (PCP) in *Drosophila* and govern ciliogenesis in vertebrates³³⁻³⁷ (Fig. 1A). We term this new module the “CPLANE,” for ciliogenesis and planar polarity effectors. We show that this module also includes the poorly understood

ciliopathy protein Jbts17, which we show recruits CPLANE to basal bodies where it acts specifically by recruiting the IFT-A peripheral proteins. In the absence of CPLANE function, the IFT-A core denuded of peripheral components still undergoes normal bi-directional transport, though the movement of IFT-B is severely impaired. Finally, examination of mutant mice and ciliopathy alleles from human patients reinforce the connection between CPLANE proteins, Jbts17, and the IFT-A machinery, demonstrating that CPLANE plays a broad and essential role in ciliogenesis and human ciliopathies.

Results:

Intu, Fuz, and Wdpcp are deeply conserved and are essential for vertebrate ciliogenesis (Fig. 1a)^{36,37}. To gain unbiased insights into their molecular functions, LAP-tagged versions of each were stably expressed in ciliated mouse kidney IMCD3 cells, lysates were prepared, and interacting proteins were affinity purified (Supplementary Fig. 1a,b). In addition, similar experiments were performed using diverse IFT-A and ciliopathy related proteins as baits. Mass-spectroscopy revealed the enrichment of the bait protein for each pull-down (Supplementary Fig. 1c, d; Supp. Dataset 1), and none of the baits were copurified with over 30 unrelated control proteins, although all baits did bring down common contaminants (Supplementary Fig. 1e). From this experiment, we identified a set of roughly 250 proteins that were individually pulled down by each of the three CPLANE proteins (Supplementary Fig. 2a).

The CPLANE interactome.

We identified an extensive interaction network for the CPLANE proteins that involved a wide array of protein machines, including dynein subunits, clathrin adaptors, and chaperonins, among others (Fig. 1b and Supplementary Fig. 2b-d). Given that CPLANE proteins are essential for vertebrate ciliogenesis^{36,37}, we were surprised to find only selective links to known ciliogenesis proteins. No links were found within the combined CPLANE interactome to many major ciliary machines, including the IFT-B complex, the BBSome, the inversin compartment, and transition

zone B9 complex. An exception was the intraflagellar transport complex A (IFT-A), as between them, pulldown for Intu, Fuz and Wdpcp identified all six subunits of the IFT-A complex (Fig. 1b, c and Supplementary Fig. 2c), though no IFT-B proteins were found. Moreover, while pulldown of IFT-A proteins themselves also efficiently returned other IFT-A proteins, pulldowns with six other ciliogenesis-related baits did not (Supp. Fig. 1d). These data suggest that CPLANE may regulate IFT-A function, which we address below.

Most strongly enriched in all CPLANE pulldowns were the CPLANE proteins themselves, with Intu, Fuz, and Wdpcp reciprocally co-purified in all combinations (Supp Figs. 1c and 2d; Supp. Table 1). By contrast, these three proteins were entirely absent from pull-downs of six additional ciliopathy related proteins (Supp. Fig. 1c). Interaction among some CPLANE proteins has also been observed in high-throughput screens of human proteins^{38,39}. Moreover, the interactions are conserved in *Drosophila*⁴⁰, though these proteins have no apparent role in ciliogenesis in that animal. In addition, the Rsg1 GTPase, which we had previously observed as a Fuz interacting protein^{38,39,41}, was also strongly associated with Intu and Wdpcp (Supp. Fig. 1c).

Finally, among the most strongly enriched proteins in the CPLANE interactomes was the largely uncharacterized ciliopathy protein Jbts17 (also called C5orf42)(Fig. 1b and Supplementary Figs. 1c and 2c). This novel protein is associated with Joubert, Oral-Facial-Digital (OFD), and Meckel Gruber syndromes⁴²⁻⁴⁵, and one recent paper identifies it as a component of the transition zone⁴⁶. We confirmed interaction between Jbts17 and all three CPLANE proteins by co-IP of *in vitro* translated proteins (Supp. Fig. 2e). Because so little is known of Jbts17, we explored its link to the CPLANE proteins in more detail.

CPLANE protein localization at the base of cilia.

Jbts17 is evolutionarily conserved (Fig. 1a) and implicated in varying ciliopathies, but its function remains largely undefined. In *Xenopus*, *Jbts17* was expressed in ciliated tissues (not shown), and knockdown using anti-sense morpholino-oligonucleotides (MOs) to disrupt splicing resulted in ciliopathy-related developmental defects, including failure of neural tube closure, defective Hedgehog signaling, and defective left/right patterning (Fig. 2a-c and Supplementary Fig. 3a,b). Accordingly, ciliogenesis was disrupted in the developing neural tube, in the node, and in multiciliated cells (MCCs)(Fig. 2d-f). CRISPR-based disruption of *Jbts17* also disrupted ciliogenesis, and co-injection of mRNA encoding GFP-Jbts17 rescued both the neural tube

defects and ciliogenesis defects resulting from MO-based knockdown (Fig. 2f and Supplementary Fig. 3a, h).

Given the physical association of Fuz, Intu, and Wdpcp with Jbts17 and Rsg1 (Supp. Fig. 1c), we explored the interrelationships among these proteins in *Xenopus* MCCs, which provide an effective platform for *in vivo* ciliary cell biology. Intu, Fuz, Rsg1, and Jbts17 localized robustly around basal bodies but were difficult to detect in axonemes (Fig. 2g). Super-resolution imaging revealed that GFP-Jbts17 was present in a ring surrounding the basal body, as marked by centrin4-BFP; this ring was similar to that formed by the distal appendage marker Cep164 (Fig. 2h). However, Jbts17 knockdown did not significantly affect recruitment of the distal appendage proteins Cep164 or Ofd1, the transition zone marker Mks1, or the basal body docking protein Hook2 (Supplementary Fig. 3g).

To assess the functional relationships of these proteins, we knocked each CPLANE protein down using MOs previously validated by mouse knockouts (see Methods) and then examined basal body localization of the remaining CPLANE members. Our data place Rsg1 at the bottom of the hierarchy: Its localization was lost from basal bodies after knockdown of any CPLANE protein, yet its own knockdown did not affect basal body recruitment of any other component (Fig. 2j and Supplementary Fig. 3c-f). Wdpcp and Intu held clear positions upstream of Fuz and Rsg1, but downstream of Jbts17. Interestingly, Wdpcp and Intu were also each required for the other's basal body localization. The role of Fuz was more complicated; unlike the other CPLANE proteins Fuz did not require Jbts17 for localization to basal bodies, and while loss of Fuz did disrupt basal body localization of Intu and Rsg1, it did not affect Wdpcp. These data provide an initial framework for the hierarchy of CPLANE protein functions.

A Jbts17 disease allele disrupts Intu localization.

Jbts17 is mutated in ciliopathies, so we asked if a disease-causing allele of Jbts17 would disrupt CPLANE function by examining a *Xenopus* cognate of a human Joubert syndrome-associated truncation of Jbts17. The R1569* truncation (equivalent to R1602 in humans⁴⁵; Supp. Fig. 4a) failed to localize to basal bodies, though another disease associated truncation R2406* localized normally, allowing us to map the basal body localization domain of Jbts17 to amino acids 1770-2318 (Supp. Fig. 4b). The pathogenicity of the R1569* truncation was apparent, because while expression of full-length Jbts17 efficiently rescues NTDs resulting from Jbts17

knockdown, expression of Jbts17-R1569* did not (Supplementary Fig. 3a). Linking this embryological defect to cell biological function, Jbts17-R1569* also failed to localize to basal bodies (Supplementary Fig. 4b) and unlike wild type, expression of Jbts17-R1569* could not rescue basal body recruitment of Intu after Jbts17 knockdown (Supplementary Fig. 4c). Thus, the Jbts17 disease allele fails to support CPLANE localization and function.

CPLANE recruits peripheral IFT-A proteins to basal bodies.

To understand the mechanisms by which CPLANE proteins impact ciliogenesis, we returned to our proteomic dataset. We were intrigued by associations between CPLANE proteins and the IFT machinery (Fig. 1b), because while we previously implicated Fuz and Rsg1 in IFT^{47,48}, their mechanism of action remained ill-defined. CPLANE proteins did not interact with IFT-B components, but rather specifically interacted with both the IFT-A core and peripheral IFT-A subunits (Fig. 1b and Supplementary Fig. 1c). Nothing is known about the regulation of peripheral sub-unit interaction with the IFT-A core, but we reasoned that by interacting with both, CPLANE could facilitate IFT-A function. Because CPLANE localizes to basal bodies, we tested this notion by assessing basal body recruitment of all six IFT-A proteins after CPLANE disruption.

Strikingly, Jbts17 knockdown specifically disrupted the recruitment of the peripheral IFT-A subunits to the basal body. The levels of Ift139, Ift121 and Ift43 at basal bodies were dramatically reduced after Jbts17 knockdown, though recruitment of the three IFT-A core proteins Ift140, Ift144, Ift122 was not disrupted (Fig. 3 and Supplementary Fig. 5a). In fact, recruitment of Ift122 was consistently increased (Fig. 3b,c). Though IFT-B is also composed of core and peripheral components, Jbts17 knockdown did not disrupt recruitment of either the peripheral IFT-B proteins Ift20, Ift80, and Cluap1, or the core IFT-B protein Ift81 (Fig. 3c and Supplementary Fig. 5b). Finally, the link between Jbts17 and IFT-A is likely relevant to disease pathology, because while full-length Jbts17 expression rescued the loss of IFT-A basal body recruitment after Jbts17 knockdown, the Joubert-associated Jbts17-R1569* truncation did not (Supplementary Fig. 4d).

Defective IFT-B trafficking in the absence of CPLANE.

IFT particles exchange rapidly between axonemes and a cytoplasmic pool around the basal body, so we were curious to know what effect failure to recruit peripheral IFT-A proteins to basal bodies after CPLANE loss would have on IFT trafficking in the axoneme. We used high-speed confocal imaging to assess IFT in MCCs and found that disruption of either *Jbts17* or *Wdpcp* disrupted IFT-B movement, as we previously found for *Fuz*⁴⁷. Kymography revealed that both peripheral and core components of IFT-B formed stationary accumulations (Fig. 4a, b; Supplementary Fig. 5f), and quantitative microscopy confirmed a significant enrichment of total IFT-B levels in the axoneme (Supp. Fig. 5h).

These accumulations are reminiscent of the effect of genetic disruption of IFT-A^{9,10}. However, the mechanisms by which peripheral IFT-A proteins impact the movement of the IFT-A core has not been examined in detail, and overall, very little is known about the dynamics of IFT-A in vertebrates. We therefore also examined the effect of CPLANE loss on the dynamics of IFT-A peripheral and core proteins in the axoneme. We found that peripheral IFT-A proteins that were not recruited to basal bodies were likewise absent from axonemes after *Jbts17* knockdown, consistent with a role for CPLANE in assembly of peripheral subunits onto the IFT-A core (Supplementary Fig. 5e). By contrast, IFT-A core proteins were not only present in axonemes at normal levels after *Jbts17* knockdown (Supp. Fig. 5d, 5h), but they also underwent bi-directional transport (Fig. 4c, d; Supp. Fig. 5g), suggesting that the core was intact and could associate with both anterograde and retrograde motors. Similar effects on IFT-A and IFT-B were observed after *Wdpcp* knockdown (Supplementary Fig. 5f,g). These data suggest that CPLANE acts by recruiting peripheral IFT-A proteins to the basal body for assembly onto the IFT-A core.

OFD Syndrome phenotypes in mouse CPLANE mutants.

Together with our proteomic data from mammalian cultured cells, the *in vivo* imaging data from *Xenopus* suggest that *Intu*, *Fuz*, and *Wdpcp* are intimately associated with *Jbts17*, which is the major gene mutated in human Oral-Facial-Digital Syndrome Type 6 (OFD6)⁴³. Key diagnostic features of OFD6 include high arched palate, tongue hamartoma, and polydactyly characterized

by Y-shaped metacarpals⁴⁹. Examining these same features in *Fuz* mutant mice, we consistently observed high arched palates, lobulated tongues and Y-shaped metacarpals (Fig. 5a-d;f, h)(see also ref. ⁵⁰). We also observed Y-shaped metacarpals and defects in tongue and palate morphology in *Wdpcp* mutants (Fig. 5e, g). These genetic data linking CPLANE components to developmental defects characteristic of OFD are consistent with our proteomic link between CPLANE and Jbts17.

Human CPLANE genes are mutated in diverse ciliopathies.

Our data argue for a functional and physical association of CPLANE proteins with Jbts17 on the one hand, and with retrograde IFT machinery on the other. We therefore examined CPLANE gene sequences in the exomes of human ciliopathy patients, focusing on OFD for its connection to Jbts17 (Ref. ⁴³) and on Short-Rib Polydactyly Syndrome (SRPS) for its association to IFT-A (Refs. ^{51,52}).

OFD VI, for which Jbts17 is the major gene⁴³, shares substantial phenotypic overlap with OFD II, including high arched palate, tongue hamartomas, and Y-shaped metacarpals^{53,54}. Consistent with our data above, exome sequencing of OFD patients with these characteristics revealed disease-associated mutations in both *WDPCP* and *INTU*. In one patient, a 5-year-old male presenting with facial dysmorphism, tongue hamartoma, high arched palate, tooth abnormalities, and postaxial polydactyly, we found trans-heterozygous mutations in *WDPCP*. One mutation was a frameshift (c.526_527delTT; Leu176Ilefs*21), and the other, a missense mutation (c.160G>A; p.Asp54Asn) predicted to alter splicing by Human Splice Finder and ASSP (Fig. 6a,b and Supplementary Table 2a). Sanger sequencing confirmed the variants and compound heterozygosity for *WDPCP* by parental segregation (Fig. 6a). The altered aspartic acid residue (D54) of *WDPCP* is highly conserved (Supplementary Fig. 6a) and has been associated with atypical OFD in one previous patient⁵⁵, but no disease associated alleles of *WDPCP* have yet been tested functionally. We therefore expressed these proteins in *Xenopus* MCCs and found that the frame-shift allele resulted in total loss of protein, while the point mutation led to a consistent but more modest defect in protein stability (Fig. 6c).

We also examined a 10-year-old male presenting with facial dysmorphisms, tongue nodular tags, high arched palate, and bilateral central Y-shaped metacarpals (Supplementary Fig. 6b and Supplementary Table 2a). Exome sequencing revealed a homozygous frameshift mutation

(c.396delT; p.Asn132Lysfs*11) in the *INTU* gene, and Sanger sequencing confirmed that the homozygous *INTU* variant segregated with the phenotype (Supplementary Fig. 6c). Brain MRI for both OFD patients showed no cerebellar abnormality, but cardiac ultrasound revealed aortic coarctation and tetralogy of Fallot in OFD patients with *CPLANE* mutations (Supplementary Table 2a), consistent with recent data implicating *CPLANE* genes in cardiac malformation⁵⁶.

In addition, we identified a homozygous missense mutation (p.Ala452Thr) in *INTU* in a child with nephronophthisis and growth retardation. The boy, who presented with end-stage renal failure at 10 yrs. of age, was the only affected child of three in a consanguineous family (Supplementary Table 2a). The identified allele segregated with the affected status, and homozygosity mapping revealed that the genomic locus of *INTU* was located within a stretch of homozygosity on chromosome 4 (Supplementary Fig. 6d and data not shown). This change is in a relatively poorly conserved residue (Supplementary Fig. 6e), suggesting that it may be a hypomorphic allele. We also identified a heterozygous mutation (p.Glu365Gly) in *WDPCP* in a boy with cerebellar vermis hypoplasia, ataxia, and retinal dystrophy (Supplementary Table 2a), which may be a modifying allele contributing to this ciliopathy.

Finally, we examined exomes of individuals with SRPS and identified an affected patient with compound heterozygosity for two *INTU* mutations confirmed by Sanger sequencing (c. 1063G>T; p.Glu355* and c. 1499A>C; p.Glu500Ala) (Fig. 6d, e; Supp. Fig. 6f). The affected individual was born pre-term at 30-weeks with multiple congenital anomalies, including wide-open fontanel, microphthalmia, tongue hamartomas, and tetralogy of Fallot (Supplementary Table 2b). Radiographic analyses showed multiple skeletal anomalies, including short horizontal ribs, shortened long bones with smooth edges, and pre-and post axial polydactyly (Fig. 6e and Supplementary Table 2b). To confirm that the identified alleles caused the patient's disease, we examined the function of the *INTU* mutants. The Glu355* allele failed to localize to basal bodies when expressed in *Xenopus* MCCs (Fig. 6f), and while the Glu500A mutation did not affect basal body localization, it did significantly impair the ability of the protein to recruit Ift43 (Fig 6g, h).

Another case, presented with strikingly similar SRPS features (Supplementary Fig. 6g,h and Supplementary Table 1b) and a single truncating mutation in *INTU* was confirmed by Sanger sequencing (c. 826C>T; p.Gln276*). This change was inherited from the unaffected mother (Figure S6i), but despite extensive analysis, no other changes were found in *INTU*. We

examined over 500 changes in the patient's genome that segregated with the phenotype, and we found only a single change in a gene previously associated with SRPS, a heterozygous change in *WDR35* (IFT121). This change (c.932G>T; p.Trp311Leu) was confirmed by Sanger sequencing and was inherited from the unaffected father (Supplementary Fig. 6i), and this residue is invariant in vertebrates (Supplementary Fig. 6j). The clinical presentation of this patient was remarkably similar to the individual with compound heterozygosity for *INTU* mutations (Supplementary Table 2b). The most striking similarity was the distinct polydactyly; while most SRPS patients have 6-7 fingers and toes, these two patients both had 9-10. Thus, while we cannot rule out a role for mutations in unknown ciliopathy genes, the known role for *WDR35* in SRPS, our finding of a role for *INTU* in SRPS, and our finding of physical and functional interactions between IFT-A and *CPLANE* lead us to suggest that SRPS in this patient results from digenic inheritance of mutations in *WDR35* and *INTU*.

Discussion:

Here, we characterize an essential but poorly understood protein module, which we term CPLANE (Ciliogenesis and Planar Polarity Effector). The CPLANE proteins Intu, Fuz, and Wdpcp/Fritz are deeply conserved in animal evolution (Fig. 1a) and were first identified as PCP proteins in *Drosophila*^{33-37,40}. In vertebrates, mutation of these genes elicits ciliogenesis defects, and while their role in PCP remains murky, recent reports do suggest that Wdpcp and Intu may govern the localization of core PCP proteins⁵⁷⁻⁵⁹. Here, we combined proteomics in mammalian cell culture, *in vivo* cell biology in *Xenopus*, and genetic analysis in both mice and humans to demonstrate that the CPLANE is an important physical and functional unit governing a specific facet of ciliogenesis, namely the recruitment of IFT-A proteins to the base of cilia and the insertion of complete IFT-A particles into the axoneme (Fig. 7a,b).

Guided by our proteomic screen, imaging experiments here revealed that in the absence of CPLANE, peripheral IFT-A proteins fail to localize to basal bodies and do not assemble onto the IFT-A core. This connection between CPLANE and IFT-A is strengthened by genetic data linking CPLANE proteins to SRPS, which is generally associated with defects in IFT-A^{51,52}. CPLANE does not traffic along axonemes, arguing that it is not a component of the IFT particle itself, a result supported by the fact that CPLANE proteins associate relatively strongly with one another in our proteomic data, but comparatively weakly with IFT proteins (Supplementary Fig. 1c). Together, these data argue that CPLANE facilitates IFT-A recruitment and assembly at basal bodies, though the precise mechanisms by which it acts remain to be determined.

One possibility is that CPLANE may direct cytoplasmic transport of IFT-A proteins to the base of cilia. This idea is supported by the many links to cytoplasmic dynein machinery in our proteomics (Fig. 1; Supp. Fig. 2). A second line of evidence comes from computational modeling of protein structures (see Methods), which suggests that CPLANE proteins contain structural domains common in vesicle trafficking machinery (Supp. Fig. 7). Fuz contains a longin domain, and Intu is predicted to fold into a Sec23/24 domain. Likewise, Wdpcp is predicted to form an alpha solenoid attached to a beta-propeller, a configuration present in coat proteins, as well as IFT and BBSome subunits⁶⁰. Finally, Jbts17 is an enormous protein (3167 amino acids) and can be threaded onto multiple protein models, including a Sec23/24 fold, a beta propeller fold, and an importin domain.

On the other hand, the CPLANE proteins may instead impact IFT-A at the level of protein stability and assembly. This possibility is supported by our proteomic data linking Intu, Fuz, and Wdpcp individually to all eight subunits of the CCT chaperone complex (Fig. 1b; Table 1). This latter

mechanism would be particularly intriguing, because similar mechanisms have been proposed linking protein folding to assembly of dynein arms^{22,23} and the BBSome^{27,28}. Importantly, as is true for the CPLANE proteins, disruption of cytosolic assembly factors for dynein arms and the BBSome also cause ciliopathies. This latter possibility is especially exciting because, despite this long history of study, cytosolic factors facilitating basal body recruitment and assembly of IFT particles have not been described.

We also found that peripheral IFT-A proteins were not present in axonemes after CPLANE knockdown, though IFT-A core proteins entered axonemes and trafficked bi-directionally. Because we also observed accumulations of IFT-B in axonemes similar to that following direct disruption of IFT-A⁹⁻¹¹, our data argue that the peripheral components are essential for association of IFT-B with IFT-A, at least during retrograde transport (Figs. 7a,b). This role for the peripheral IFT-A proteins raises interesting questions concerning the mechanism of IFT-B anterograde transport, because biochemical data suggest that heterotrimeric kinesin also associates with IFT particles via the IFT-A^{7,13}. Given that IFT-B moves into axonemes and accumulates there after CPLANE knockdown (Supp. Fig. 5c), it may be that the role for peripheral proteins in linking IFT-A to IFT-B is specific to retrograde transport, consistent with the known remodeling of IFT particles upon turnaround^{11,12}. Alternatively, homodimeric kinesin 2 associates directly with IFT-B in *C. elegans*, and while the situation in vertebrates is less clear^{61,62}, IFT-B could move into cilia after CPLANE knockdown via direct association with the homodimeric Kif17. Finally, the role for diffusion in axonemal transport is only now coming into focus^{20,63}, so it is at least possible that IFT-B in this cases moves in the axoneme could move in such a manner.

Finally, our data implicate CPLANE in diverse ciliopathies. Because SRPS is generally associated with disruption of IFT-A^{51,52}, our identification of *INTU* mutations in SRPS patients provides a genetic complement to our proteomic and cell biological linkage of CPLANE to IFT-A. Our identification of CPLANE mutations in OFD patients similarly parallels the physical and functional association of CPLANE with Jbts17. Additional insights into the mechanism of CPLANE mutation in human disease will certainly emerge as we venture deeper into the CPLANE protein interaction network. For example, *Wdpcp* mutations have been found in at least one patient with Bardet- Biedl Syndrome³⁶, so the links between CPLANE proteins and CCT are interesting since BBS6, 10 and 12 also interact with CCT^{27,28}. Our data may also shed light on the still murky role of *FUZ* in human neural tube defects⁶⁴. Finally, our data suggest that the CPLANE may be relevant not only to congenital defects, but also to infectious disease, since recent data link *Fuz* and the endocytic machinery to alphavirus entry⁶⁵, and we observed extensive interactions between

498 CPLANE and the vesicle trafficking machinery (Fig. 1b; Supp. Fig. 2b). In sum, our data
499 establish the CPLANE as a novel ciliogenic protein module with significant roles in
500 development and human disease.

Figure legends

Figure 1. The CPLANE interactome. (a) CPLANE proteins are deeply conserved in evolution; numbers indicate percent identity (+additional percent similarity). “XX” indicates that reciprocal BLAST identified no clear orthologue. (b) The extended CPLANE protein network. Tandem affinity purification of Intu, Fuz, Wdpcp, IFT-A¹⁵, and the published NPHP network⁴ were assembled, thresholded for most likely network members, and over 2200 nodes arranged in Cytoscape, with 78 nodes presented here. The raw peptide data for this network will be deposited in the appropriate databases upon acceptance of the manuscript.

Figure 2. Jbts17 localizes to the base of cilia and is required for ciliogenesis and cilia-mediated patterning. (a) RT-PCR demonstrates disrupted Jbts17 splicing after MO injection. ODC1 = loading control. (b) *In situ* hybridization of SHH direct target, *nkx2.2* in Jbts17-knockdown (st. 22). (c) *Pitx2* expression at st. 26. Arrows indicate signal in left lateral plate mesoderm (LPM); graph indicates *pitx2* expressing embryos. (d) Acetylated α -tubulin immunostaining in the ventral neural tube (st. 22)(Scale bars, 10 μ m). (e) Nodal cilia length is reduced; cilia numbers are unchanged. (Scale bars, 10 μ m). Graphs in D and E each show pooled data from two independent experiments cilia length (mean \pm SEM; ***p <0.001). (f) Multi-ciliated cells in control, Jbts17-knockdown (Jbts17-KD), and rescue with untargeted Jbts17 mRNA. Cilia and cell membrane visualized by GFP-CFAP20 (green) and membrane-RFP (magenta). Scale bars, 10 μ m. (g) GFP-tagged Jbts17 localizes near basal bodies (visualized by co-expressed centrin4-RFP) in an MCC (scale bar, 10 μ m). (h) Super-resolution image of GFP-Jbts17 and mCherry-Cep164 at a single basal body; both form rings of \sim 260nm around the basal body, visualized by centrin4-BFP. Diameters shown as mean \pm SD in each panel. The lower graph shows fluorescence intensities for GFP-Jbts17, mCherry-Cep164, and centrin4-BFP (Scale bar, 100 nm). (i) GFP-tagged CPLANE proteins (green) and basal bodies visualized by centrin4-RFP (magenta) in control and Jbts17 knockdown multi-ciliated cells; box plots of CPLANE intensities at basal bodies; boxes extend from 25th-75th percentiles, with a line at the median; whiskers indicate max and min. (j) Table summarizes the localization of CPLANE proteins at basal bodies for each knockdown (see Supplementary Fig. 2c-f).

Figure 3. Jbts17 is necessary for recruitment of peripheral IFT-A proteins to basal bodies. (a) Ift43 localization at basal bodies in *Xenopus* MCCs as marked by centrin4-RFP is lost in MCCs after Jbts17 knockdown. (b) Peripheral IFT-A components are not recruited to centrin4-RFP labeled basal bodies after Jbts17 knockdown. (c) Quantification of IFT protein localization to basal bodies from two independent experiments. Graphs show fluorescence intensity of GFP fusions to indicated IFT proteins normalized against that of centrin4-RFP (see methods). Peripheral IFT-A proteins are specifically lost after Jbts17 knockdown.

Figure 4. Jbts17 is required for bi-directional axonemal transport of IFT-B particles, but not the core IFT-A. Still images from high-speed time-lapse movies of IFT using GFP fusion to IFT proteins (green) and membrane-RFP (magenta). Cluap1-GFP in control embryo (a) and Jbts17 morphant (b). GFP-IFT144 in control (c) and Jbts17 morphant (d). Insets show high magnification views of localization of IFT particles in a single axoneme in the boxed regions.

Scale bars = 10 μ m. Associated kymograph representing movements of IFT particles are shown in panels a'-d'.

Figure 5. CPLANE mutant mice display diagnostic features of Oral-Facial-Digital Syndrome Type 6. (a and b) Frontal sections at E14.5 reveal that *Fuz* mutant mice display high arched palate (arrow) and lobulation of the tongue (arrowheads). (c and d) *Fuz* mutant mice display develop polydactyly with Y-shaped metacarpals. (e) *Wdpcp* mutant mice display develop polydactyly with Y-shaped metacarpals. (f, g, h) Frontal sections of E13.5 Wildtype, *Wdpcp* and *Fuz* mutant embryos. DAPI labels nuclei (cyan). (f', g', h') Illustrations highlighting the corresponding palatal condensations (purple) and tongue (pink) in F-H. Mutant palatal condensations form more medially than do controls and fail to extend into the mouth (See Tabler et al., 2013).

Figure 6. CPLANE mutations in human ciliopathies. (a) Pedigree showing *WDPCP* mutations in an OFD patient. (b) The patient displays tongue hamartomas and dental anomalies. (c) When expressed in *Xenopus* embryos, the D54A allele of human *WDPCP* produces less protein compared to wild-type; the L176F-fs26* allele produces no protein. (d) Pedigree showing *INTU* mutations in an SRPS phenotype. (e) X-ray of the patient. (f) The E355* allele of *INTU* disrupts basal body localization. (g and h) Wild-type *Intu* rescues *Ift43* localization to basal bodies after *Intu* knockdown; the E500A allele of *Intu* does not. Data shown are pooled from three independent experiments.

Figure 7. Models for CPLANE function and structure. (a) Schematic of normal IFT. Peripheral proteins are assembled onto the IFT-A core in the cytoplasm and injected together with IFT-B for bi-direction transport in axonemes. (b) In the absence of CPLANE, IFT-A core particles lacking peripheral proteins are injected into axonemes and traffic normally; IFT-B enters axonemes but fails to move in a retrograde direction and accumulates.

Experimental procedures:

Details of the experimental procedures can be found in the Supplementary Material online. Briefly, the methods used here were essentially those described previously. Immunoprecipitation of LAP-Tagged CPLANE proteins was performed as described in ref. ⁴. General methods for experiments with *Xenopus* were performed as described in ref ^{36,37}. Specifically, imaging of IFT proteins at basal bodies and time-lapse imaging of IFT were performed as per Ref. ⁴⁷. Exome sequencing was performed essentially as per Ref. ⁴ and Ref. ⁴³.

Online Supplementary Experimental Procedures:

Tandem Affinity Purification and Mass Spectrometry

To best optimize the LAP purification procedure and minimize the possibility of carryover, we have standardized the growth of cells, the preparation of extracts, and the method of tandem affinity purification (Torres et al., 2009). Briefly, stable LAP cell lines were harvested using detergent. The mIMCD-3 (ATCC® CRL-2123™) cell line was obtained from ATCC. The renal IMCD3 line was authenticated by expression of renal markers (AQP2), polarization, establishment of tight junctions, and ciliation, as outlined in (REF). Lysates were clarified at 43,000 rpm and subjected to

anti-GFP immunoprecipitation. Bound proteins were eluted from antibody beads using TEV protease, recaptured on S-protein agarose (Novagen), and eluted in 4x NuPAGE sample buffer (Invitrogen). Following purification, great care is taken to ensure a lack of contamination from both environmental sources and from other purified proteins. Each purified set of interacting proteins is separated on an individual 10% Bis-Tris polyacrylamide gel and stained with Coomassie brilliant blue. IMCD3 samples were run into gels for 20-40mm and divided into 20-40 x1 mm slices. Each excised lane was reduced, carboxyamidomethylated and digested with trypsin. Peptide identification of each digestion mixture was performed by microcapillary reversed-phase HPLC nanoelectrospray tandem mass spectrometry (mLC-MS/MS) on an LTQ-Orbitrap Velos or XL mass spectrometer (ThermoFisher Scientific, San Jose, MA). The Orbitrap repetitively surveyed an m/z range from 395 to 1600, while data-dependent MS/MS spectra on the twenty (Velos) or ten (XL) most abundant ions in each survey scan were acquired in the linear ion trap. MS/MS spectra were acquired with relative collision energy of 30%, 2.5-Da isolation width, and recurring ions dynamically excluded for 60 s. Preliminary sequencing of peptides was facilitated with the SEQUEST algorithm with a 30 ppm mass tolerance against a species specific (mouse or human) subset of the UniProt Knowledgebase. With a custom version of the Harvard Proteomics Browser Suite (ThermoFisher Scientific, San Jose CA), peptide spectrum matches (PSMs) were accepted with mass error < 2.5ppm and score thresholds to attain an estimated false discovery rate (FDR) of <1% using a reverse decoy database strategy. For IMCD3-LAP-GFP control samples, peptide identification was performed using NanoAcquity UPLC (Waters Corp, Milford, MA). Samples were analyzed on-line via nanospray ionization into a hybrid LTQ-Orbitrap mass spectrometer (Thermo Scientific, San Jose, CA). Data were collected in data dependent mode with the parent ion being analyzed in the FTMS and the top 8 most abundant ions being selected for fragmentation and analysis in the LTQ. Tandem mass spectrometric data were analyzed using the Mascot search algorithm (Matrix Sciences, London, UK) against the mouse Uniprot database (including reverse hits and contaminants) and filtered to a 5% FDR.

TAP/MS data network generation

For individual genes identified in each AP/MS sample, we calculated the normalized spectral abundance factor (NSAF)

$$NSAF = \frac{P_g/L_g}{\sum_i P_i/L_i}$$

Where P_g represents the number of spectral counts mapped to gene g , and L_g is the mean length, in residues, of reference protein isoforms derived from that gene. The P/L quotient is normalized to the sum of P/L quotients for all genes in the dataset, excluding those derived from the bait protein and those derived from known exogenous proteins. We exclude, for example, proteins commonly found in skin and enzymes added during sample preparation. Using an unpublished set of eight negative control datasets, we systematically search for genes whose score in an experimental data set is highly unlikely. These filtered genes included in the attached Cytoscape (<http://www.cytoscape.org>) network file, and a manually curated, simplified subset of these are shown in Figure 1b.

Xenopus embryo manipulations and microinjections

Female adult *Xenopus laevis* were induced to ovulation by injection of human chorionic gonadotropin, and eggs were fertilized *in vitro*, dejellied in 3% cysteine (pH 7.9) at 2-cell stage,

and subsequently reared in 0.3× Marc's Modified Ringer's (MMR) solution. For microinjections, embryos were placed in a solution of 2% Ficoll in 0.3× MMR and handled using forceps and an Oxford universal micromanipulator. After injections, embryos were reared for 3 hr in 2% Ficoll in 0.3× MMR and then washed and reared in 0.3× MMR until the appropriate stages. All experiments were repeated in multiple embryos from multiple clutches, as is standard in the field. No explicit randomization or blinding strategy was employed. Sex cannot be practically determined for tadpoles used in these studies. This work was carried out with approval of the University of Texas Animal Care and Use Committee.

Morpholinos, mRNA, and gRNA injections

Capped mRNA was synthesized using mMESSAGE mMACHINE kit (Ambion). mRNA and antisense morpholino were injected into two ventral blastomeres or two dorsal blastomeres at the 4-cell stage to target the epidermis or the neural tissues, respectively⁶⁶. Morpholino sequences and the working concentrations include:

We note that the *Intu*, *Fuz*, and *Wdpcp* MO used here elicit phenotypes validated by genetic disruption of these genes in mice^{36,37,57,67}. The *Rsg1* MO has been validated by disruption of the *Rsg1* GTPase by expression of a dominant-negative mutant⁴¹.

Plasmid Constructs

Three cDNA fragments encoding *Xenopus laevis* *Jbts17* were amplified by PCR from stage 25 *Xenopus* cDNA library with the primers

These three PCR products were then fused to generate full-length cDNA by PCR with *Sma*I and *Not*I restriction enzyme sites at 5 and 3 prime ends, respectively. Truncated mutants were generated by PCR. The cDNAs were then subcloned into various pCS107 vectors that established fusion with either EGFP, mRFP, or myc-tag at the N-terminus.

Full-length *Odf1*, *Cep164*, *Hook2*, *Mks1*, *IFT122*, *IFT140*, *IFT144*, *IFT43*, *IFT121*, *IFT139*, *IFT81*, *IFT80* and *Cluap1* cDNAs were identified in *Xenbase*⁶⁸ and amplified by PCR and sub cloned into pCS107 vectors fused with GFP, RFP, or mCherry at their N-termini. *Rsg1*, *IFT20*, *IFT43*, *Centrin4* and *Inturned* constructs were previously described in^{47,48,69}. GFP-tagged CFAP20 was kindly gifted from Dr. Brian Mitchell (Northwestern Univ.).

In vitro protein translation and pull down assay

GFP-tagged *Inturned*, *Fuzzy*, *WDPCP* and Flag-tagged *Jbts17* (907-1569) proteins were synthesized using a TnT® SP6 Coupled Wheat Germ Extract System (Promega). After centrifugation at 17,000 *g* at 4°C for 10 minutes, the synthesized soluble proteins were mixed and incubated in binding buffer (20 mM Tris-HCl pH 8.0, 1 mM EDTA, 1 mM DTT and 150 mM NaCl) for 2 hr at 4°C. Flag-tagged *Jbts17* protein was precipitated by anti-Flag M2 magnetic beads (Sigma-Aldrich). The beads were washed with binding buffer containing 0.1% Triton X-100 four times and eluted with 200 ng/ml of Flag peptide in 1x TBS. The eluted proteins were analyzed by immunoblotting with anti-GFP (Abcam) and anti-Flag M2 (Sigma-Aldrich) antibodies.

sgRNA synthesis, Cas9 induced genomic editing and genotyping

Cas9-CRISPR mediated genome editing in *Xenopus* was previously described (Nakayama et al., Methods Enzymol 546:355-75 (2014)).

oligonucleotides. The amplicon was purified by PCR purification kit (Qiagen) and used as a template for sgRNA synthesis. sgRNA was generated by using T7 MEGAscript kit (Ambion) and purified by MEGAclean Transcription Clean-up kit (Ambion). 1.25 ng of Cas9 protein (PNABIO) and 250 pg of sgRNA were injected into animal pole at 1-cell stage. Genomic DNA was extracted from stage 25 embryos using DNeasy Blood & Tissue kit (Qiagen) following manufacturer's protocol and the 468 bp of genomic region which contains the sgRNA target sequence was amplified by PCR.

The genome editing efficiency by CRISPR was examined by T7 Endonuclease I (T7EI) assay and NcoI digestion and analyzed by 2% agarose gel.

Basal body imaging and quantification

Fluorescence images of GFP, RFP or BFP fusion proteins at basal bodies in stage 25-28 multiciliated cells were captured with LSM700 inverted confocal microscope (Carl Zeiss) with a Plan-APOCHROMAT 63×/1.4 oil immersion objective.

The fluorescence intensities of GFP, RFP, or BFP fused proteins at basal bodies in slices from 2 mm of apical surface was measured by the 3D object counter plug-in of Fiji software as described previously^{47,48}. The Object size was set to 20, and threshold was determined empirically to maximize detection of apparent foci. At least 14 basal bodies/cells from at least 12 cells in 4 independent embryos were analyzed. Statistical analysis was subsequently performed, making use of the Student t-test.

Live imaging of IFT particles and analysis

For high-speed *in vivo* imaging, *Xenopus* embryos injected with GFP-tagged IFT mRNAs and membrane RNA were anaesthetized with 0.005 % benzocaine at stage 25-28. Images were acquired on Nikon eclipse Ti confocal microscope with a 63×/1.4 oil immersion objective. Time-lapse series were captured 266.5 msec per frame. Kymographs were generated and IFT velocities measured in Fiji with the kymograph plugin.

Super-resolution microscopy

GFP-Jbts17, mCherry-Cep164 and Centrin4-BFP were co-expressed by mRNA injection epidermis in stage 27 *Xenopus* embryos. Fluorescence signals were captured by LSM 710/Elyra S.1 (Carl Zeiss) via a 63×/1.3 oil-immersion objective. Raw fluorescence images were processed by structured illumination algorithm using Zen software (Carl Zeiss). The arbitrary unit (a.u.) of fluorescence intensities of ring-shaped structure of GFP-Jbts17 and mCherry-Cep164 at basal bodies were measured by Plot Profile plugin in Fiji software.

Immunostaining

Immunostaining was performed as per⁷⁰. Embryos were fixed in MEMFA for 2 hr at room temperature after removing the vitelline envelop and washing with PBST (1 ×PBS containing 0.05% TritonX-100). Transverse sections of neural tube were generated using a Vibratome series 1000 from stage 22 embryos, and GRP region was manually dissected from stage 17 embryos using forceps. Sections were dehydrated completely in methanol at −20 °C overnight. After rehydration, sections were blocked with 10% fetal bovine serum in PBST for 60 min, incubated with anti-acetylated- α -tubulin (1:2000, clone 6-11B-1, Sigma), and diluted in blocking solution. Primary antibody was conjugated with an Alexa Fluor 488 goat anti-mouse IgG (Invitrogen). Fluorescence images were obtained using Zeiss LSM700 confocal microscope. Cilia length in neural tube and GRP was measured in Fiji software.

***In situ* hybridization**

Whole mount *in situ* hybridization of *Xenopus* embryos was performed as described previously⁷¹ using the Dig-labelled RNA probe against *nkx2.2*, *vax1* and *pitx2c*. Bright field images were captured on a Zeiss Axiozoom V16 stereomicroscope.

RT-PCR

Jbts17 morpholino was injected into the all cells at 4-cell stage, and the total RNA was isolated using TRIzol reagent (Invitrogen) from four stage 25 *Xenopus* embryos. The stage 25 cDNA library was synthesized using M-MLV Reverse Transcriptase (Invitrogen) with random hexamers. Jbts17 and an Ornithine decarboxylase 1 (ODC1) internal control were amplified using Phusion® High-Fidelity DNA Polymerase (New England Biolabs) with following primers:

The abundance of the resulting amplicon was analyzed on a 2.0% agarose gel stained with ethidium bromide under UV transilluminator.

Protein structure modelling

For fold recognition studies of Fuz, Intu, Wdpcp, Jbts17 and Rsg1, we used HHpred⁷² and pGenThreader⁷³ with default parameters searched against the latest set of PDB structures. We used the respective Modeller pipelines on the HHpred and pGenThreader servers to build homology models based on high confident fold recognition results. Validity of the homology models was checked using automated scores provided by HHpred. Additionally, DaliLite⁷⁴ was used to align the homology models with the original template for visual inspection of the fold.

Human subjects

The committees approving work with human subjects include the GAD collection n° DC2011-1332 (Thauvin-Robinet), Institutional Review Boards of the University of Michigan and Boston Children's Hospital (Hildebrandt), and the Human Subjects Institutional Review Board of the University of California at Los Angeles (Krakow). Informed consent was obtained from all study participants and consent was obtained for use of all images of human subjects.

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Author Contributions:

Michinori Toriyama contributed to design, execution and interpretation of the overall research plan, with special emphasis on all frog embryo experiments and in vitro binding assays. He also contributed to writing the manuscript. Chanjae Lee designed, performed and interpreted live-imaging of IFT particles in axonemes and contributed to other imaging experiments in *Xenopus*. Kevin Drew and Edward M. Marcotte provided protein structural models. Jacqueline M. Tabler, Jiang Chen and Karen J. Liu contributed to design execution and interpretation of mouse genetic data. Marcus R. Kelley contributed to the execution and analysis of the proteomic data. Tae Joo Park contributed to *Xenopus* studies. S. Paige Taylor, Ivan Duran, Daniel H. Cohn, Ange-Line Bruel, Daniella Braun, Ghislaine Pierquin, Armand Biver, Kerstin Wagner, Anne Malfroot, Inusha Panigrahi, Brunella Franco, Hadeel Adel Al-lami, Yvonne Yeung, Yeon Ja Choi, University of Washington Center for Mendelian Genomics, Yannis Duffourd, Laurence Faivre, Jean-Baptiste Rivière contributed to collection of human patient and sequencing data. Friedhelm Hildebrandt, Christel Thauvin-Robinet, and Deborah Krakow contributed to design, execution and interpretation of human genetic data. Peter K. Jackson contributed to coordinating the overall research effort with a focus on design and interpretation of the proteomic screen and contributed to writing the manuscript. John Wallingford coordinated the overall research effort, oversaw experimental design and interpretation, and wrote the manuscript.

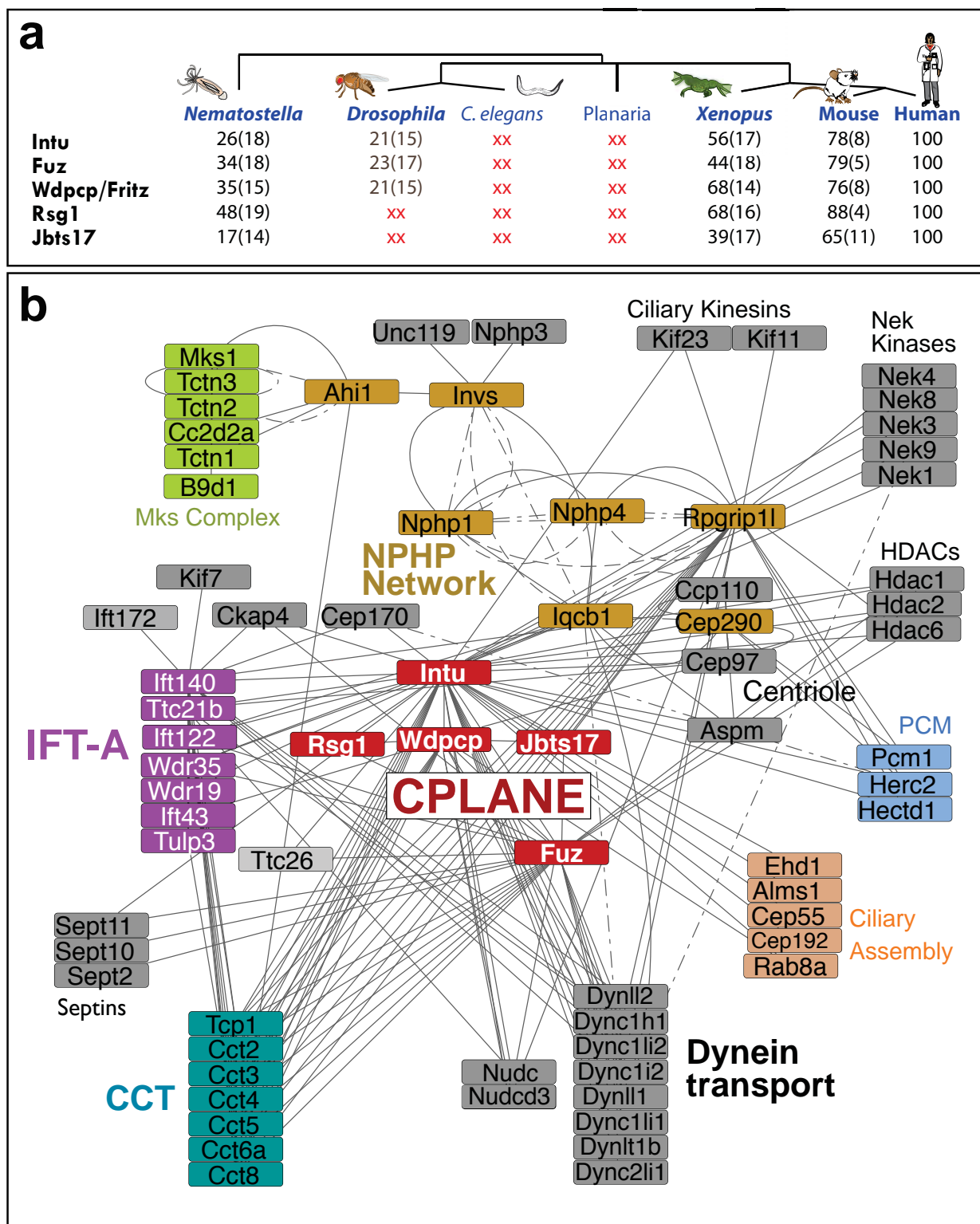
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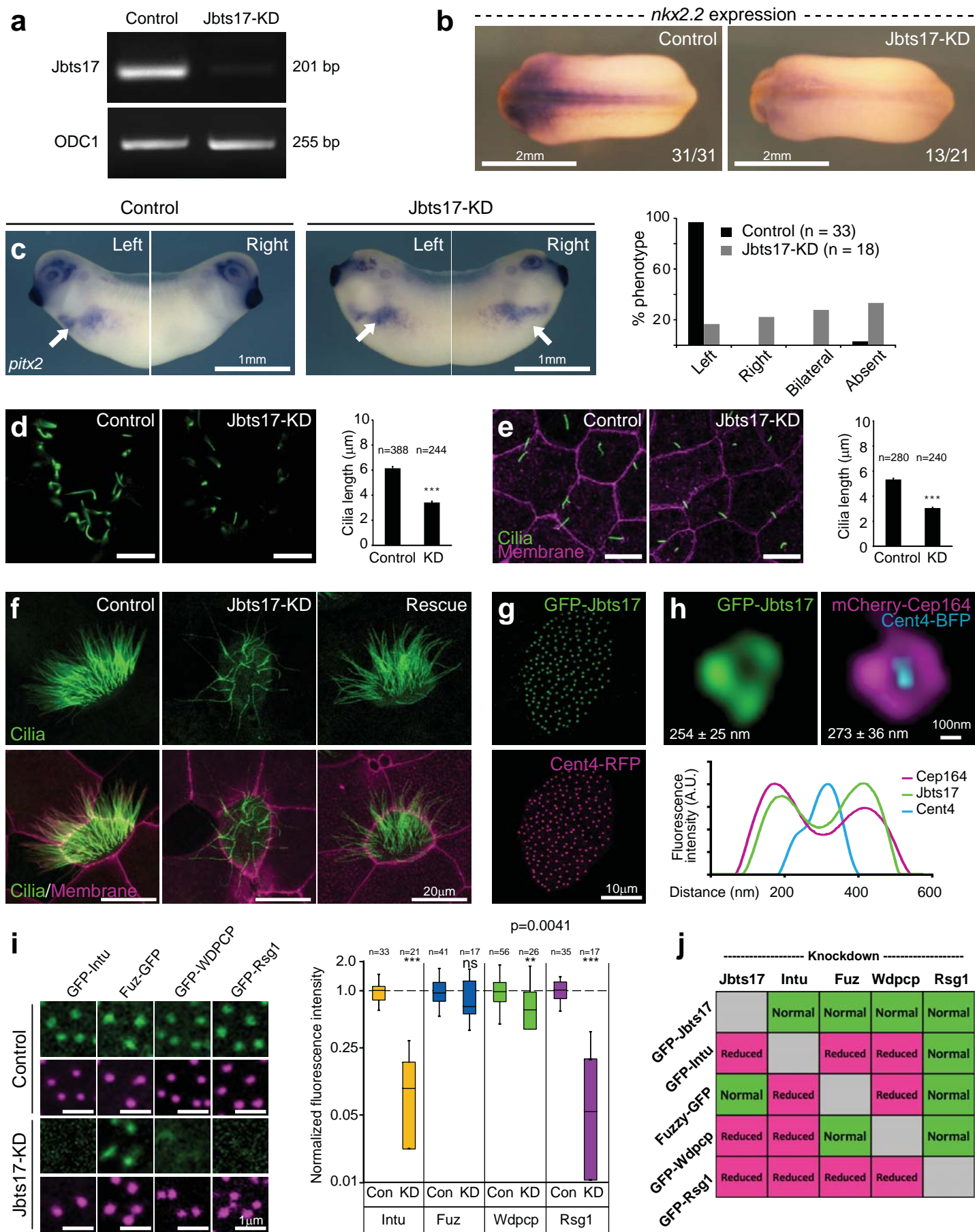
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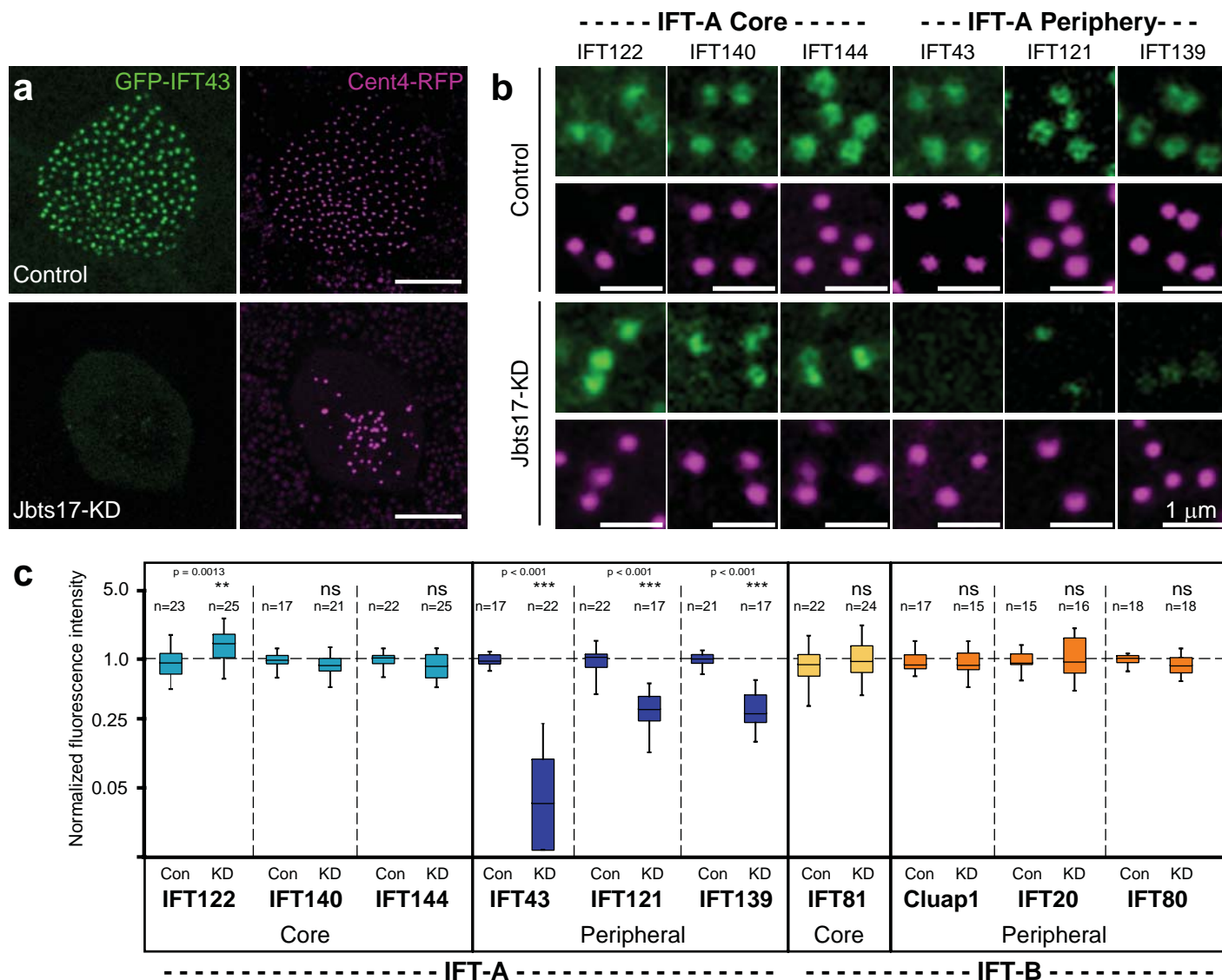
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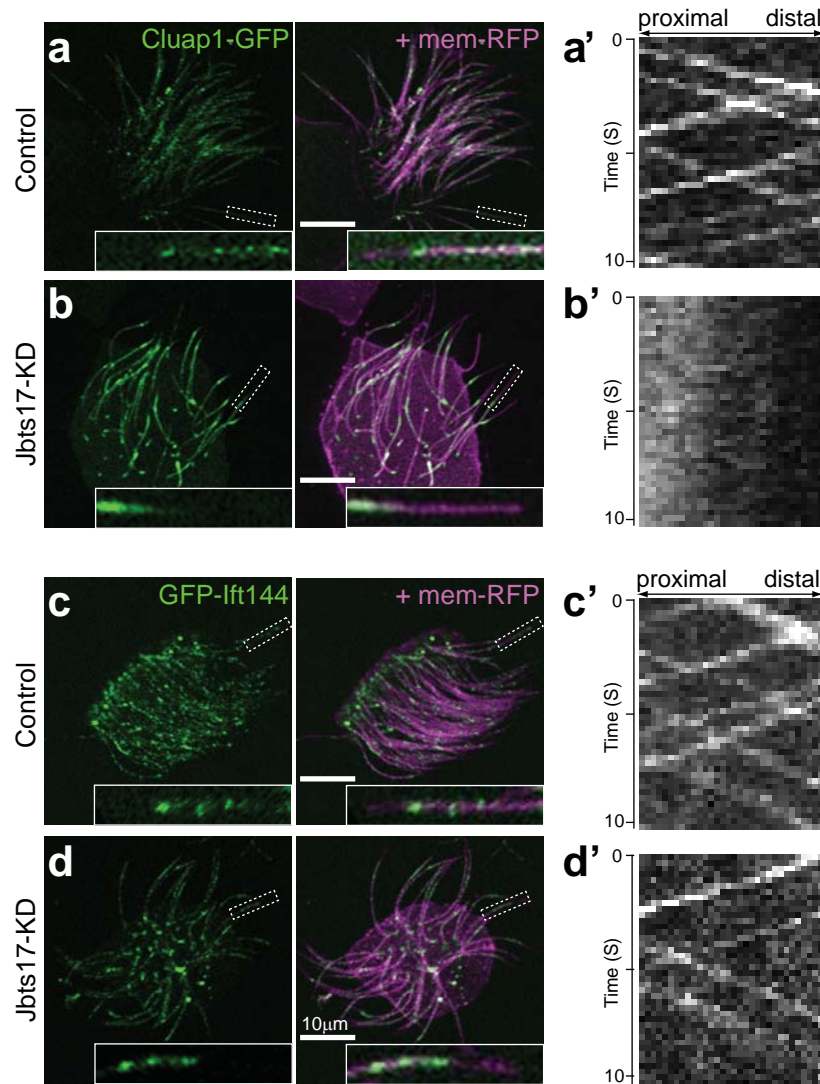
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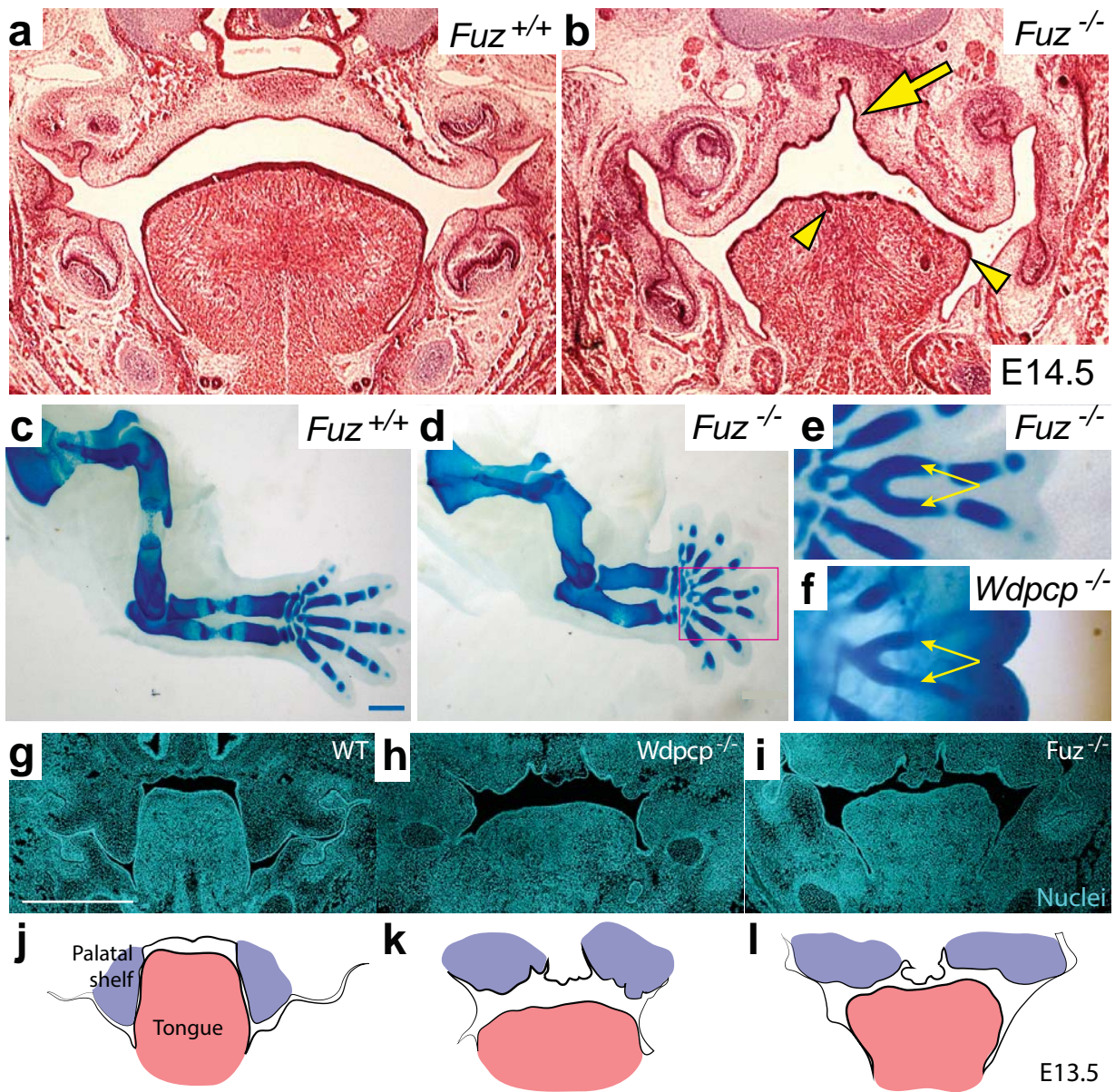




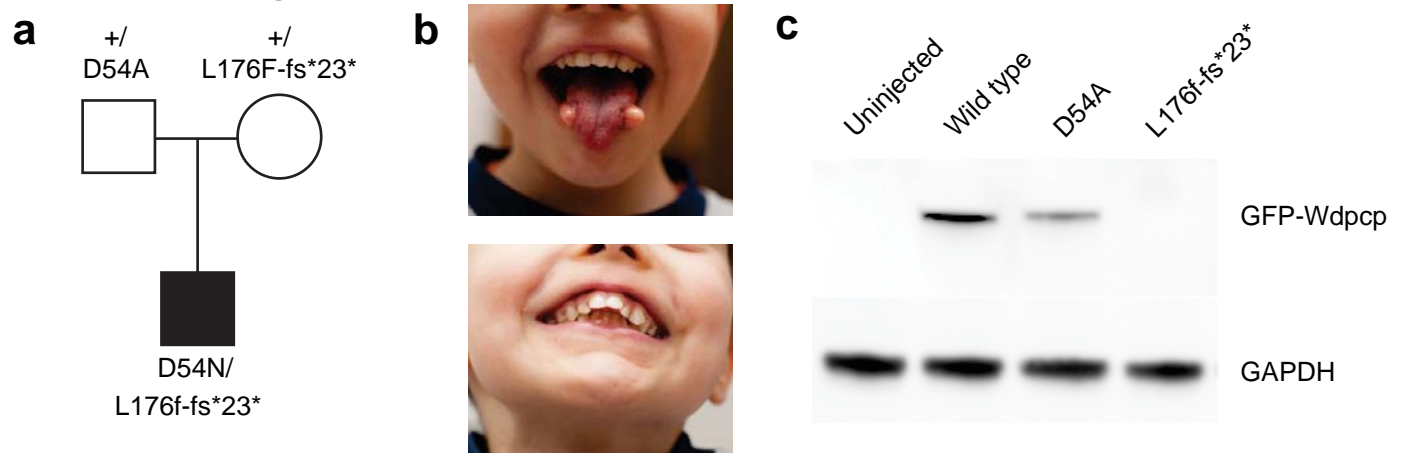




Toriyama et al. Figure 4



Oral-Facial-Digital - *WDPCP*



Short Rib Poydactyly - *INTU*

